

Glucocorticoid receptor–JNK interaction mediates inhibition of the JNK pathway by glucocorticoids

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Inhibition of the c-Jun N-terminal kinase (JNK) pathway by glucocorticoids (GCs) results in AP-1 repression. GC antagonism of AP-1 relies mainly on the transrepression function of the GC receptor (GR) and mediates essential physiological and pharmacological actions. Here we show that GCs induce the disassembly of JNK from mitogen-activated protein kinase kinase 7 (MKK7) by promoting its association with GR. Moreover, we have characterized a hormone-regulated JNK docking site in the GR ligand-binding domain that mediates GR–JNK interaction. The binding of GR to JNK is required for inhibition of JNK activation and induction of inactive JNK nuclear transfer by GCs. The dissociation of these two hormone actions shows that JNK nuclear transfer is dispensable for the downregulation of JNK activation by GCs. Nonetheless, nuclear accumulation of inactive JNK may still be relevant for enhancing the repression of AP-1 activity by GCs. In this regard, chromatin immunoprecipitation assays show that GC-induced GR–JNK association correlates with an increase in the loading of inactive JNK on the AP-1-bound response elements of the *c-jun* gene.

Keywords: AP-1 antagonism/cell signaling/cross-talk/ MAPK docking site/MAPK pathway

Introduction

Glucocorticoids (GCs) play key physiological roles in development, cellular proliferation and differentiation. In addition, the prominent pharmacological actions of these hormones have prompted their widespread medical use to treat diverse pathological conditions such as asthma, allergic rhinitis, rheumatoid arthritis and leukemia (Barnes, 1998). GCs exert most of their actions by binding to an intracellular GC receptor (GR), a ligand-activated transcriptional regulator that belongs to the nuclear receptor (NR) superfamily (Beato *et al.*, 1995).

In most circumstances, hormone-free GR is associated with heterotypic complexes that contain chaperones, such as Hsp90, and co-chaperones, and is retained in the cytoplasm. Upon ligand binding, the chaperone complex is released and hormone-bound GR is rapidly transferred into the nucleus. Hormone-activated GR regulates gene transcription, either positively or negatively, by two major modes of action. The most well known involves the binding of GR homodimers to the GC response elements (GREs) found in the regulatory sequences of GC target genes. A second and more elusive mode of action is independent of the direct interaction of GR with DNA and relies on the interference (thus, also known as cross-talk or transrepression) with the activity of other transcriptional regulators by mechanisms based on protein–protein interactions. In contrast to the former, transrepression is apparently mediated by GR monomers. In fact, transactivation-defective mutants of GR, which cannot dimerize (GR^{dim}) or bind DNA (GRLS7), are fully competent in transrepression (Heck *et al.*, 1994; Helmsberg *et al.*, 1995). Remarkably, the *in vivo* relevance of the DNA binding-independent actions of GR has been evidenced by the generation of mice that harbor the GR^{dim} mutation (Reichardt *et al.*, 1998). Unlike the GR-deficient mice, which show severe abnormalities and die shortly after birth (Cole *et al.*, 1995), GR^{dim} mice are viable, indicating that the DNA binding-dependent activities of GR are dispensable for survival (Reichardt *et al.*, 1998). Important sets of genes transrepressed by GR are those which are under the positive control of AP-1 and/or NF- κ B. As these transcriptional regulators play critical roles in controlling the expression of many proinflammatory genes, GR antagonism with AP-1 and NF- κ B is believed to underlie the anti-inflammatory and immune-suppressive actions of GCs (Göttlicher *et al.*, 1998; Herrlich, 2001). The maintenance of GC anti-inflammatory activity together with the ability to transrepress AP-1 and NF- κ B in GR^{dim} mice indicate that this pharmacological action is mostly mediated by the transrepression function of the GR (Reichardt *et al.*, 2001).

The antagonism between GR and AP-1 was described in the early 1990s, and several underlying mechanisms have since been proposed (reviewed in Herrlich, 2001). While early studies suggested a mutual inhibition to bind to DNA because of the formation of a GR–AP-1 complex, genomic footprinting (Konig *et al.*, 1992) and chromatin immunoprecipitation (ChIP) assays (Rogatsky *et al.*, 2001) showed that GC repression occurs with promoter-bound AP-1, suggesting that interaction with GR prevents AP-1 from proper interactions with the transcriptional machinery or with a co-activator complex (Saatioglu *et al.*, 1994) and/or recruits co-repressor complexes (Rogatsky *et al.*, 2001).

An alternative mechanism by which GCs may exert their antagonistic action on AP-1 is through GR-mediated

interference of the signaling pathways that activate AP-1, in particular the mitogen-activated protein kinase (MAPK) pathways (Caelles *et al.*, 2002). MAPKs contribute to AP-1 induction in response to a series of extracellular

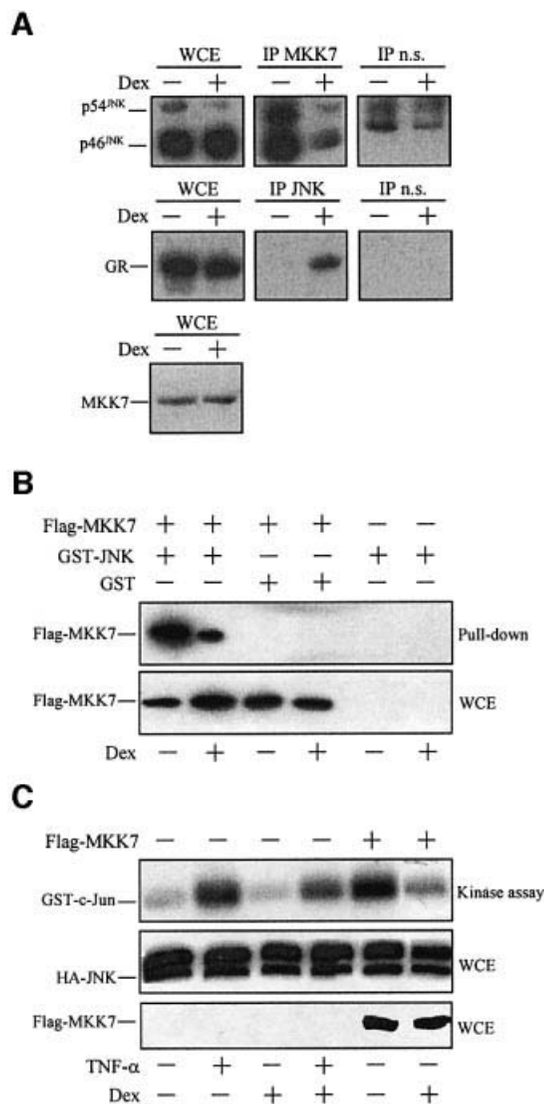


Fig. 1. GCs induce reduction of the MKK7-JNK complexes and promote GR-JNK association. (A) Extracts from serum-starved HeLa cells incubated with Dex (or vehicle) were immunoprecipitated with antibodies to MKK7 (IP MKK7), JNK (IP JNK) and, as a non-specific antibody, GST (IP n.s.), respectively. Immunoblots were performed to analyze the presence of JNK and GR in the immunocomplexes and the JNK, GR and MKK7 content in the whole-cell extracts (WCE), as indicated. (B) HeLa cells were transiently co-transfected with either pCM15-Flag-MKK7 or pCM15 along with pEBG-JNK or pEBG, serum starved and treated with Dex (or vehicle). GST or GST-JNK was precipitated from cell extracts by glutathione-Sepharose beads and the associated Flag-MKK7 was examined by immunoblotting using an anti-Flag antibody (upper panel). The lower panel shows the protein level of Flag-MKK7 in the cell extracts assessed by immunoblotting using an anti-Flag antibody. (C) HeLa cells were transiently co-transfected with pCEFL-KZ-HA-JNK together with pCM15-Flag-MKK7 or empty vector, serum starved and treated with Dex (or vehicle). When indicated, extracts were prepared after stimulation with TNF- α . The activity of HA-JNK was determined by immunocomplex assay (upper panel) and the amount of HA-JNK (middle panel) and Flag-MKK7 (lower panel) by immunoblotting using anti-HA and anti-Flag antibodies, respectively.

stimuli. Interestingly, AP-1 induction by distinct MAPKs is usually mediated by phosphorylation of a particular set of substrates and therefore involves distinct mechanisms (Karin, 1995). In recent years, compelling evidence has highlighted the relevance of MAPK pathways as targets of GC action. Indeed, GC-induced repression of AP-1 activation may be achieved by GR-mediated inhibition of the c-Jun N-terminal kinase (JNK) pathway. This GC action leads to the inhibition of phosphorylation, and concomitant activation, of JNK-targeted transcriptional activators, such as c-Jun, ATF-2 or Elk-1, which are involved in the induction of AP-1 activity by different mechanisms (Caelles *et al.*, 1997). Other MAPK pathways, such as the extracellular signal-regulated protein kinase (ERK) and p38 MAPK cascades, are also targets for repression by GCs (Caelles *et al.*, 2002). GCs may interfere with these signaling pathways through the expression of the dual-specificity MAPK phosphatase-1 (MKP-1). This mechanism has been shown to account for the inhibition of ERK (Kassel *et al.*, 2001) and p38 MAPK pathways (Imasato *et al.*, 2002; Lasa *et al.*, 2002) in a number of cell types. Although MKP-1 induction may also be involved in GC-induced inhibition of the JNK pathway, alternative mechanisms should exist since GCs can inhibit JNK activation even in the absence of *de novo* gene expression (Caelles *et al.*, 1997; Ventura *et al.*, 1999).

Here we report that GR physically interacts with JNK through a hormone-regulated JNK docking site located in the ligand-binding domain (LBD) of GR. This GC action mediates the inhibition of JNK pathway activation and the induction of inactive JNK nuclear transfer. Remarkably, both hormone effects may separately contribute to AP-1 transrepression. GR-JNK interaction constitutes a novel mechanism by which GCs regulate cellular signaling and gene expression. Additionally, as this GC action targets the AP-1 complex, this mechanism may also conduct some of the pharmacological actions of GCs.

Results

GCs reduce the number of MKK7-JNK complexes

In HeLa cells, a short GC treatment inhibits the JNK pathway and c-Jun N-terminal phosphorylation in a GR-dependent manner (Caelles *et al.*, 1997; González *et al.*, 2000; Caelles *et al.*, 2002). In parallel, this same GC treatment induces inactive JNK to accumulate in the nucleus (González *et al.*, 2000). In the light of these observations, we proposed that this latter hormone action may be involved in the interference of GCs with activation of the JNK pathway and the AP-1 complex (Caelles *et al.*, 2002). To study the molecular mechanism that underlies this interference further, we analyzed the effect of GC treatment on the integrity of mitogen-activated protein kinase 7 (MKK7)-JNK complexes by co-immunoprecipitation assays of endogenous proteins performed in extracts from HeLa cells. In agreement with previous reports, we found that JNK was associated with the MKK7 immunocomplexes in unstimulated cells (Tournier *et al.*, 1999). Remarkably, treatment with dexamethasone (Dex) clearly reduced the amount of JNK bound to MKK7 without altering the overall amount of JNK or MKK7 in the whole-cell extracts (Figure 1A, upper and lower panels, respectively). These results were corroborated

further by GST pull-down assays performed in extracts from HeLa cells transiently co-transfected with expression vectors for Flag-MKK7 and/or GST-JNK or, as a negative control, GST (Figure 1B). Moreover, the decrease in the MKK7-JNK complexes induced by Dex correlated with its inhibitory action on the JNK activity triggered by tumor necrosis factor (TNF)- α stimulation or MKK7 over-expression (Figure 1C).

GR binds to JNK through a hormone-regulated JNK docking site

The results above, together with those previously reported for GC-induced nuclear transfer of inactive JNK (González *et al.*, 2000), suggested that GR might physically interact with JNK in response to hormone. Additionally, this hypothesis was supported by previous data that showed that the kinetics of GC inhibitory action on the JNK pathway correlated with the nuclear entry of the GR (Caelles *et al.*, 1997; Ventura *et al.*, 1999). To

pursue this idea further, we analyzed GR-JNK interaction in response to Dex treatment by co-immunoprecipitation assays of endogenous proteins using extracts from HeLa cells. In contrast to the GC action on the MKK7-JNK complexes, GR co-immunoprecipitated with JNK specifically in extracts from GC-treated cells (Figure 1A, middle panels). Unfortunately, in these same assays, we did not detect JNK in GR immunoprecipitates (not shown), which may be due to the interference of the GR antibody with the GR-JNK complex by either masking the interaction surface or promoting an inadequate conformation of GR. Hormone-induced GR-JNK interaction could also be evidenced by GST pull-down assays performed with extracts from cells overexpressing GST-JNK (see below).

Most MAPK-interacting proteins contain discrete amino acid motifs that function as MAPK docking sites (Jacobs *et al.*, 1999). Thus, we searched the primary sequence of GR for potential MAPK docking sites. We found that the GR LBD contains an amino acid sequence that significantly resembles a D-box, a MAPK docking site which is characterized by a basic and a hydrophobic-X-hydrophobic submotif separated by a short spacer and found in other JNK-interacting proteins, such as c-Jun and JIP-1 (Hibi *et al.*, 1993; Dickens *et al.*, 1997) (Figure 2A). Indeed, an 11 residue peptide corresponding to the JIP amino acid sequence shown in Figure 2A is sufficient for binding to and inhibiting JNK (Barr *et al.*, 2002). Consistent with these observations, a truncated version of GR, GR-LBD, comprising 25 amino acids of the hinge region plus the whole LBD, effectively binds JNK when transiently expressed *in vivo* (Figure 2B). Moreover, this interaction is competed specifically *in vivo* by the JNKII

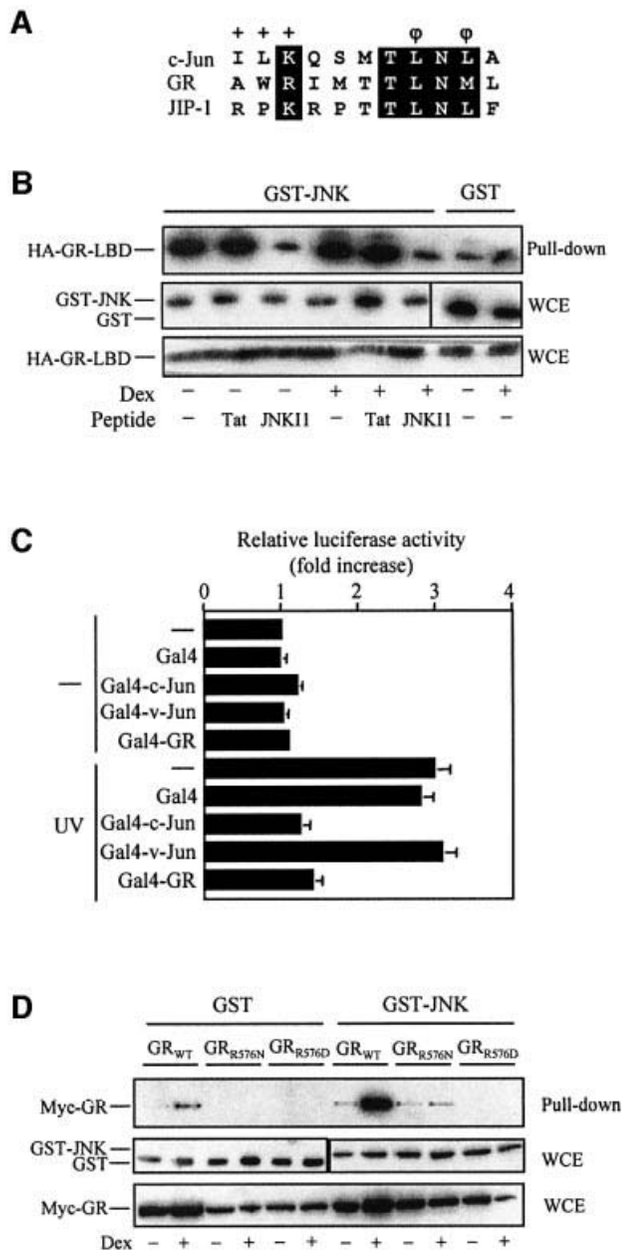


Fig. 2. GR harbors a hormone-regulated docking site for JNK. (A) Alignment of amino acid sequences of rat GR (574–584) with the JNK docking site found in human c-Jun (33–43) and JIP-1 (157–167). The characteristic features of the D-box MAPK docking site which are a basic (+) and a hydrophobic (ϕ)-X-hydrophobic submotif separated by a short spacer are indicated. Identical or conserved residues among all sequences are boxed in black. (B) Cos-7 cells were transiently co-transfected with pCEFL-KZ-HA-GR-LBD along with pEBG-JNK or pEBG, serum starved and treated with Dex (or vehicle). Thereafter, vehicle, Tat or JNKII peptides (1 μ M) were added as indicated and cells further incubated for 30 min. Cell extracts were prepared, glutathione-Sepharose precipitated and the presence of HA-GR-LBD associated with GST-JNK or GST was analyzed by immunoblotting using an anti-HA antibody (upper panel). The protein level of GST-JNK or GST (middle panels) and HA-GR-LBD (lower panel) in cell extracts was analyzed by immunoblotting using anti-GST and anti-HA antibodies, respectively. (C) HeLa cells were transiently co-transfected with the -73col-luciferase reporter (3 μ g) along with pSG424, pSG424-Gal4-c-Jun(1–116), pSG424-Gal4-v-Jun(1–89) or pSG424-Gal4-GR(540–738) (2 μ g), as indicated, and pCHI10 (0.5 μ g). After serum starvation, cells were UV stimulated, as indicated, and luciferase and β -galactosidase activities were measured after 8 h. Relative luciferase activity corresponds to the β -galactosidase-normalized luciferase activity. The fold increase compared with the activity in non-irradiated cells transfected with the reporter construct alone, which was set to 1, is shown (mean \pm SD of three independent experiments run in triplicate). (D) Cos-7 cells were transiently co-transfected with pEBG-JNK or pEBG along with pMTG-myc-GR, pMTG-myc-GR_{R576N} or pMTG-myc-GR_{R576D}, serum starved and treated with Dex (or vehicle) as indicated. Thereafter, GST pull-down assays were performed and the GST precipitates were analyzed by immunoblotting using an anti-myc antibody (upper panel). The protein levels in the whole-cell extracts of GST-JNK or GST (middle panels) and myc-GR (lower panel) were determined by immunoblotting using anti-GST or anti-myc antibodies, respectively.

peptide, a cell-permeable peptide which contains the JNK docking site of JIP $-1_{157-176}$ covalently linked to the

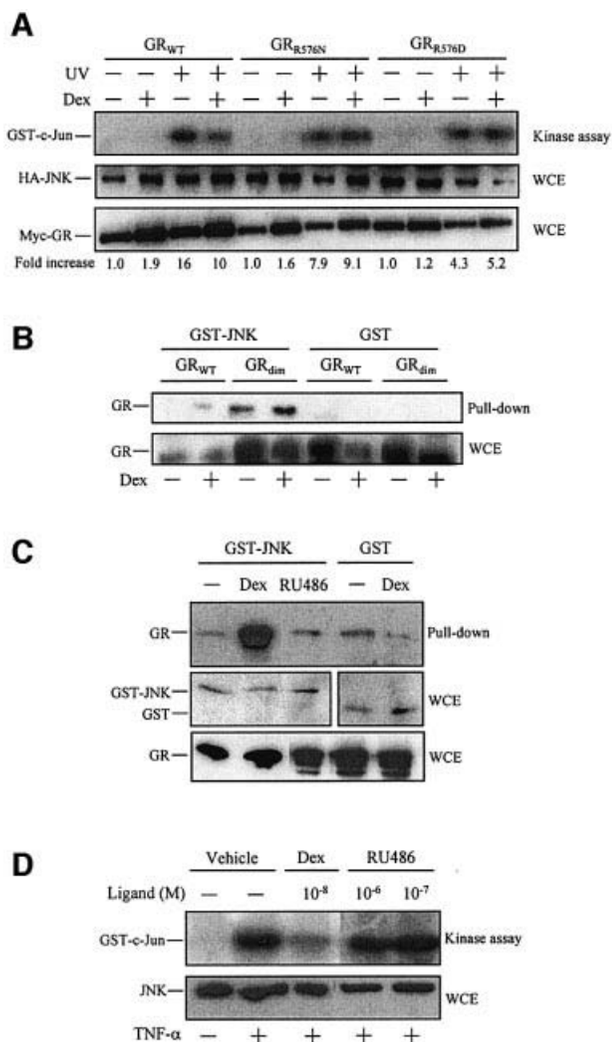


Fig. 3. GR binding to JNK mediates downregulation of JNK activation by GCs and does not require GR dimerization. (A) Cos-7 cells were transiently co-transfected with pCEFL-KZ-HA-JNK along with pMTG-myc-GR, pMTG-myc-GR^{R576N} or pMTG-myc-GR^{R576D}, serum starved, pre-treated with Dex (or vehicle) and stimulated with UV, as indicated. Thereafter, cells were harvested and the activity of HA-JNK determined (upper panel). The expression level of HA-JNK (middle panel) and myc-GR (lower panel) in each cell extract was analyzed by immunoblotting using anti-HA and anti-myc antibodies, respectively. The numbers below each lane indicate the fold increase of HA-JNK activity relative to the untreated/unstimulated condition. (B) Cos-7 cells were transiently co-transfected with pSB-GR or pSB-GR(A458T) along with pEBG-JNK or pEBG, serum starved and treated with Dex (or vehicle), as indicated. Thereafter, GST pull-down assays were performed and the GST precipitates (upper panel) and cell extracts (lower panel) were analyzed by immunoblotting using an antibody to GR. (C) HeLa cells were transiently transfected with pEBG-JNK or pEBG, serum starved and treated with Dex (10^{-8} M), RU486 (10^{-6} M) or vehicle, as indicated. Thereafter, GST pull-down assays were performed and the presence of GR in glutathione-Sepharose precipitates was analyzed by immunoblotting using an antibody to GR. The protein level in the cell extracts of GST-JNK or GST (middle panels) and GR (lower panel) was measured by immunoblotting, using antibodies to GST or GR, respectively. (D) Serum-starved HeLa cells were pre-treated with Dex, RU486 or vehicle at the doses indicated and then stimulated with TNF- α . Cell extracts were tested for JNK activity by immunocomplex assay (upper panel) and JNK protein level by immunoblotting (lower panel) using an antibody to JNK.

10 amino acid human immunodeficiency virus (HIV)-Tat₄₈₋₅₇ sequence, which acts as a carrier peptide (Bonny *et al.*, 2001). Interestingly, although the GR-LBD still exhibits ligand modulation in relation to JNK interaction to some extent, it binds JNK even in the absence of hormone (Figure 2B). Although we do not have a conclusive explanation for the partial loss of hormone regulation of this truncated GR protein, co-immunoprecipitation assays comparing the amount of Hsp90 associated with the full-length GR versus the GR-LBD showed that the latter associates with Hsp90 very poorly (data not shown). This result suggests that conformational differences between these two GR versions might exist and/or could support the hypothesis that Hsp90 might be somehow involved in masking the JNK docking site.

Additionally, we tested the ability of a region of GR that encompasses this putative JNK docking site to repress JNK pathway-dependent transcription as evidence that this interaction takes place in living cells. As shown in Figure 2C, a Gal4-GR(540-738) fusion protein inhibits the JNK-mediated activation of an AP-1-dependent reporter as efficiently as Gal4-c-Jun(1-116), which contains the JNK docking site of c-Jun. In contrast, Gal4-v-Jun(1-89), which is defective in the JNK-docking site and, hence, in interaction with JNK, or the Gal4 DNA-binding domain alone have no effect. None of these constructs had any significant effect on the basal expression of the AP-1-dependent reporter in unstimulated cells.

Notably, the amino acid sequence of this putative JNK docking site of GR is not fully conserved in any other member of the NR superfamily, not even in its closest relative, the mineralocorticoid receptor (MR) in which several non-conservative amino acid changes affect, respectively, the basic and the hydrophobic-X-hydrophobic submotifs. In particular, the basic motif of GR is not conserved in either human or murine MR due to the substitution of Arg576 by an asparagine or serine, respectively. Since it has been shown that the basic submotif is critical for JNK binding (Ho *et al.*, 2003), to abrogate GR interaction with JNK we mutated Arg576 of GR to asparagine (GR^{R576N}) or aspartate (GR^{R576D}). At the Dex concentration tested (10^{-6} M), both GR mutants trigger transcription to the same level as the wild-type receptor, as shown by transient transfection assays using a 2xGRE-luciferase reporter (data not shown). In contrast, GST pull-down assays showed that both GR mutants are defective in hormone-induced binding to JNK (Figure 2D).

GR-JNK interaction correlates with inhibition of JNK activation

Next, we used several approaches to study whether GC-induced inhibition of JNK correlates with JNK binding to GR. First, we tested whether the JNK interaction-defective mutants of GR, GR^{R576N} and GR^{R576D}, failed to mediate GC-induced inhibition of JNK activation. As shown in Figure 3A, in contrast to the wild-type receptor, both JNK interaction-defective mutants of GR are also defective in JNK inhibition.

Previously, we have shown that the dimerization-defective GR mutant GR^{dim} is as efficient as the wild-type receptor in mediating downregulation of JNK activity by GCs (González *et al.*, 2000; Caelles *et al.*, 2002). Therefore, here we tested the GR^{dim} mutant for

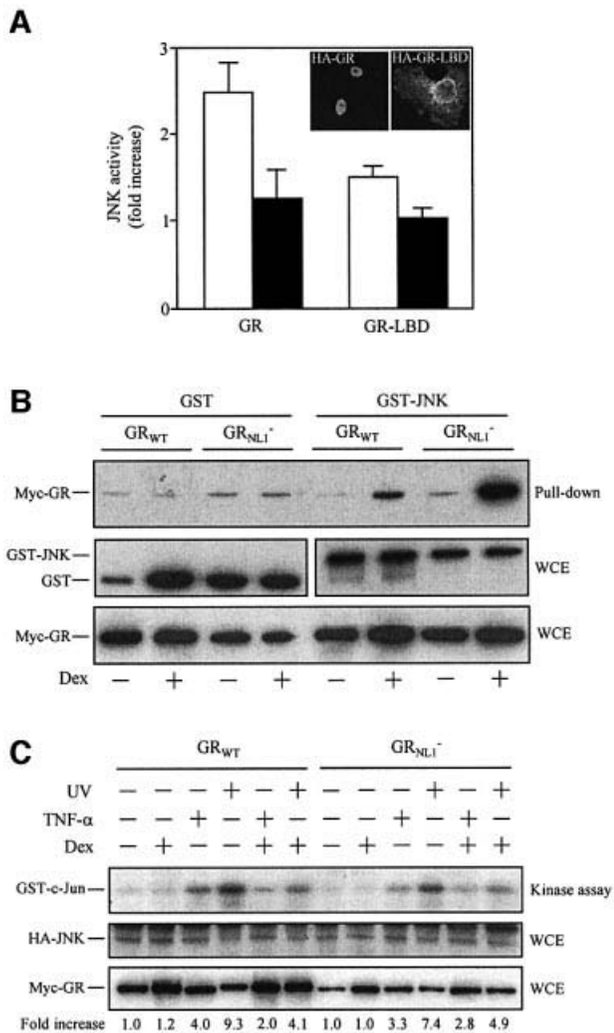


Fig. 4. Hormone-induced nuclear transfer of GR is dispensable for binding to and downregulation of JNK. **(A)** Cos-7 cells were transiently co-transfected with pCEFL-KZ-HA-JNK together with pEBG-GR or pEBG-GR-LBD, serum starved and treated with Dex (or vehicle) for 45 min before TNF- α stimulation. Cell extracts were prepared 20 min after stimulation and the HA-JNK activity determined by immunocomplex assay. The graphic shows the increase of JNK activity in response to TNF- α stimulation in the absence (white bars) or presence (black bars) of Dex. The average results from three independent experiments are shown. The inset shows an anti-HA immunocytochemical analysis by confocal microscopy of Dex-treated Cos-7 cells transiently transfected with pCEFL-KZ-HA-GR or pCEFL-KZ-HA-GR-LBD, as indicated. **(B)** Cos-7 cells were transiently co-transfected with pMTG-myc-GR or pMTG-myc-GR_{NLI}⁻ along with pEBG or pEBG-JNK, serum starved and treated with Dex (or vehicle), as indicated. GST pull-down assays were performed and precipitates analyzed for the presence of myc-GR by immunoblotting with an anti-myc antibody (upper panel). The GST or GST-JNK and myc-GR protein level in each cell extract was analyzed by immunoblotting using specific anti-GST or anti-myc antibodies, respectively (middle and lower panel, respectively). **(C)** Cos-7 cells were transiently co-transfected with pCEFL-KZ-HA-JNK along with pMTG-myc-GR or pMTG-myc-GR_{NLI}⁻, serum starved, pre-treated with Dex (or vehicle) and stimulated with UV or TNF- α , as indicated. Thereafter, cells were harvested and the activity of HA-JNK determined (upper panel). The expression level of HA-JNK (middle panel) and myc-GR (lower panel) in each cell extract was analyzed by immunoblotting using anti-HA and anti-myc antibodies, respectively. The numbers below each lane indicate the fold increase of HA-JNK activity relative to the untreated/unstimulated condition.

hormone-induced binding to JNK. For this purpose, GST pull-down assays were performed in extracts from Cos-7 cells overexpressing GST-JNK or, as a negative control, GST together with wild-type GR (GR_{WT}) or GR^{dim}. As shown in Figure 3B, GR^{dim} is fully competent at mediating GC-induced binding to JNK.

Additionally, we also compared the ability of the GR agonist Dex versus the antagonist RU486 to induce binding to and inhibition of JNK. Unlike Dex, RU486 failed to induce GR to either interact with or inhibit activation of JNK (Figure 3C and D, respectively). Altogether, these results strongly support the correlation between GR-JNK interaction and inhibition of JNK activation by GCs. Furthermore, the results obtained with GR^{dim} indicate that GR dimerization is dispensable for GR binding to JNK.

GC-induced nuclear transfer of JNK is mediated by binding to GR but is dispensable for inhibition of JNK activation by GCs

The GR-LBD truncated protein, which is fully competent in binding to JNK, lacks the nuclear localization signal 1 (NL1). Consistently, GR-LBD is not translocated efficiently into the nucleus upon ligand activation, as shown by immunocytochemistry analysis (inset in Figure 4A), even though expression of the GR-LBD efficiently inhibits the TNF- α -induced activity of JNK when overexpressed in Cos-7 cells (Figure 4A). In fact, similarly to the binding to JNK, the GR-LBD inhibits JNK activation even in the absence of hormone. Hormone addition only weakly, though significantly, improved JNK inhibition by this truncated form of GR. On the basis of this observation, we examined whether these two hormone actions, inhibition of JNK activation and induction of inactive JNK nuclear transfer, could be dissociated in the context of the full-length receptor. For this purpose, we took advantage of the GR_{NLI}⁻ mutant, which, because of three point mutations in the NL1, is highly defective in hormone-induced transfer into the nucleus (Savory *et al.*, 1999). First, the GR_{NLI}⁻ mutant was tested for its ability to interact with JNK. Ligand-induced binding of GR_{NLI}⁻ to JNK was evidenced by GST pull-down assays performed with extracts from Cos-7 cells overexpressing either GR_{WT} or GR_{NLI}⁻ along with GST-JNK or GST (Figure 4B). We next compared GC action on the UV- and TNF- α -induced activation of JNK mediated by either wild-type or NL1-defective GR. Hormone-activated GR_{NLI}⁻ was as efficient as the wild-type receptor in inhibiting the activation of JNK (Figure 4C). In a manner similar to GR-LBD, this mutated version of GR, engineered to remain in the cytoplasmic compartment upon hormone activation, can mediate inhibition of the JNK pathway.

We also used this NL1-defective GR mutant to assess whether JNK binding to GR dictates JNK subcellular trafficking in response to GCs. Immunofluorescence analysis by confocal microscopy showed that while the GR_{WT} translocates into the nucleus and induces nuclear accumulation of JNK in response to Dex, GR_{NLI}⁻ did not mediate either of these two hormone actions (Figure 5A). To support this argument further, we took advantage of the JNK interaction-defective mutants GR_{R576N} and GR_{R576D}, as well as the fact that RU486 is fully competent in inducing GR nuclear translocation (Htun *et al.*, 1996)

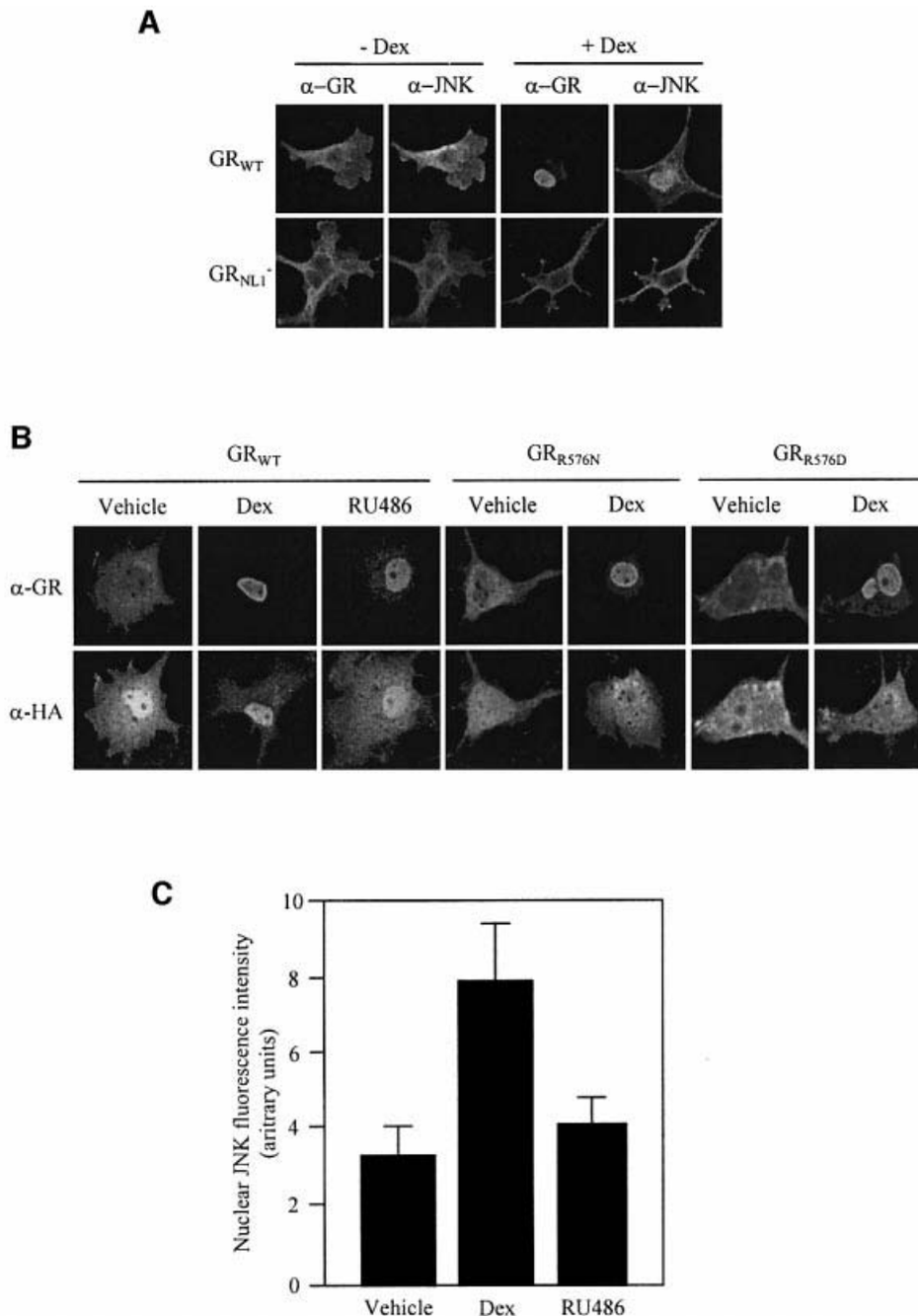


Fig. 5. JNK binding to GR mediates GC-induced nuclear transfer of JNK. **(A)** Immunofluorescence analysis by confocal microscopy of JNK and GR in Cos-7 cells transiently transfected with pMTG-myc-GR or pMTG-myc-GR_{NLI}. After transfection, cells were serum starved and fixed after treatment with Dex (or vehicle). Cells were double immunostained by using primary antibodies to JNK and GR, as described in Materials and methods. **(B)** Cos-7 cells were transiently co-transfected with pMTG-myc-GR, pMTG-myc-GR_{R576N} or pMTG-myc-GR_{R576D} along with pCEFL-KZ-HA-JNK, serum starved and treated with Dex (10^{-8} M), RU486 (10^{-7} M) or vehicle, as indicated. Thereafter, immunocytochemical analysis was performed by double staining with anti-GR and anti-HA antibodies and confocal microscopy. **(C)** Serum-starved HeLa cells were treated with Dex (10^{-8} M), RU486 (10^{-7} M) or vehicle, as indicated, and analyzed by immunocytochemistry using antibodies to GR and JNK. Immunofluorescence was analyzed by confocal microscopy. For each condition, at least 300 nuclei were analyzed and quantified as described in Materials and methods.

whereas it did not induce either binding to or inhibition of JNK. We performed immunocytochemical analysis by confocal microscopy of GR and JNK in Cos-7 cells overexpressing hemagglutinin (HA)-JNK along with GR_{WT}, GR_{R576N} or GR_{R576D}, respectively, and treated with vehicle, Dex or, when relevant, RU486. GR_{WT} as well as both GR mutants translocate into the nuclear compartment upon Dex addition. However, unlike GR_{WT},

both JNK interaction-defective mutants failed to induce the accumulation of JNK inside the nucleus (Figure 5B). In relation to the GR antagonist RU486, it induced nuclear transfer of GR which was comparable with Dex, but failed to stimulate JNK nuclear accumulation (Figure 5B). Similar results were obtained by immunocytochemical analysis of the endogenous GR and JNK proteins in HeLa cells (Figure 5C).

INPUT			IP JNK			IP n. s.		
-	D	RU	-	D	RU	-	D	RU
3'-c-jun			3'-c-jun			3'-c-jun		
5'-c-jun			5'-c-jun			5'-c-jun		
Fold increase			3.0	4.0	2.1	1.0	1.0	1.1

INPUT			IP GR			IP n. s.		
-	D	RU	-	D	RU	-	D	RU
3'-c-jun			3'-c-jun			3'-c-jun		
5'-c-jun			5'-c-jun			5'-c-jun		
Fold increase			1.8	7.5	4.2	1.0	0.9	1.1

Fig. 6. GC-induced nuclear transfer of JNK increases JNK associated with the AP-1 response elements of the *c-jun* gene. Serum-starved HeLa cells were treated with Dex (D), RU486 (R) or vehicle (-), as indicated. Thereafter, cells were processed for ChIP assays using antibodies to JNK (IP JNK), GR (IP GR) or, as a non-specific antibody (IP n.s.), GST, and primers pairs that amplify DNA fragments containing either the distal and proximal AP-1 sites of the *c-jun* promoter region (5'-c-jun) or, as a measure of PCR amplification due to non-specific immunoprecipitation, a region corresponding to the 3' end of the *c-jun* gene (3'-c-jun). Equal amounts of total genomic DNA (INPUT) were used for immunoprecipitation in each condition. For each sample, the radioactive labeling of the 5'-jun and 3'-c-jun PCR products was quantified and expressed as a 5'-c-jun:3'-c-jun ratio. The ratio obtained with GST antibody in untreated cells was arbitrarily set as 1 (values shown below the gels).

These results support our initial hypothesis that inhibition of the JNK pathway by GCs is likely to occur at the cytoplasmic level and strongly indicate that GC-induced dissociation of JNK from the MAPK module, by binding to the GR, suffices to inhibit the activation of JNK. Conversely, JNK nuclear translocation, while also being mediated by binding to GR, may be dispensable for this particular inhibitory action.

GC-induced nuclear transfer of JNK increases loading of JNK on the AP-1-bound response elements of the *c-jun* gene

Although JNK nuclear transfer does not appear to be essential for GC-induced downregulation of JNK activation, we reasoned that it could still be relevant for AP-1 transrepression. Indeed, in a previous study, we proposed that inactive JNK accumulated in the nucleus in response to GCs may act as an inhibitor of AP-1 activity by binding to the c-Jun transactivation domain and, consequently, blocking further interaction with activated JNK (González *et al.*, 2000; Caelles *et al.*, 2002). To gain evidence to support this idea, we performed ChIP assays to monitor the JNK associated *in vivo*, and in response to GCs, with the AP-1 response elements of the *c-jun* gene. In these assays, we amplify a DNA fragment (5'-c-jun) containing both the proximal and distal AP-1 response elements of the *c-jun* gene (Angel *et al.*, 1988). Importantly, both AP-1 sites mediate transrepression of the *c-jun* gene by GCs (Wei *et al.*, 1998). ChIP assays using antibodies to c-Jun showed that AP-1 complexes containing c-Jun are bound *in vivo* to the AP-1 sites of the *c-jun* gene in both non- and GC-treated HeLa cells (not shown). In agreement with previous reports showing that c-Jun and JNK

co-immunoprecipitate in resting cells (Dai *et al.*, 1995), we also found JNK specifically associated with this region in unstimulated cells. Remarkably, Dex treatment produces a modest but reproducible enrichment of JNK associated with the AP-1 response elements of the *c-jun* gene (Figure 6). ChIP assays also showed that GR is tethered to the AP-1 sites of the *c-jun* gene in response to Dex or, to a lesser extent, RU486, a result which is consistent with a previous study on the AP-1 response element of the collagenase 3 gene (Rogatsky *et al.*, 2001). However, the most striking difference between the GR agonist (Dex) and antagonist (RU486) is the efficiency of the former in tethering JNK to the AP-1 sites (Figure 6). Significantly, this divergence draws a parallel with the differences observed between these GC analogs in inducing the binding of GR to JNK.

Discussion

In recent years, several studies have described the negative regulation of MAPK pathways by GCs and have proposed its involvement in carrying out some of the physiological and pharmacological actions of these hormones. Here we show that GCs reduce the amount of JNK associated with the MAPK module by promoting its binding to the GR. Consistently, we have identified a hormone-regulated JNK docking site located within the GR LBD. We also show that GR binding to JNK mediates the inhibition of the JNK pathway and the induction of JNK nuclear transfer in response to GCs. Although JNK binding to GR is sufficient for the former GC action, we provide evidence that the latter may synergize in AP-1 transrepression by inducing the loading of inactive JNK onto the AP-1-bound response elements (see model in Figure 7).

Induction of MKP-1 expression contributes to the downregulation of the ERK and p38 MAPK pathways by GCs (Kassel *et al.*, 2001; Imasato *et al.*, 2002; Lasa *et al.*, 2002). While this mechanism may also be suitable to mediate GC-induced inhibition of the JNK pathway, alternative modes of action should exist since the downregulation of this pathway by GCs proceeds even in the presence of actinomycin D and can be mediated by transactivation-defective mutants of GR such as GR^{dim} or GRLS7 (Caelles *et al.*, 1997; Ventura *et al.*, 1999; González *et al.*, 2000). In this regard, our results support the hypothesis of an alternative mechanism based on protein-protein interactions.

We had reported previously that the step along the JNK pathway targeted by GCs is situated downstream from the MAP3K level (Caelles *et al.*, 1997). Our present data are consistent with this finding since GCs reduce the overall amount of MKK7-JNK complexes, and hence inhibit MKK7-induced activation of JNK. Moreover, we show that GC interference with the MKK7-JNK complexes correlates with the induction of GR to bind to JNK by these hormones. In summary, our results support a novel mechanism responsible for the GC inhibitory action on the JNK cascade: the disruption of the pathway by the GR-mediated sequestration of JNK. This mechanism may affect all JNK signaling pathways since this MAPK would be unavailable not only to MKK7 but also to its other MAP2K, SEK1. In this regard, we already showed that

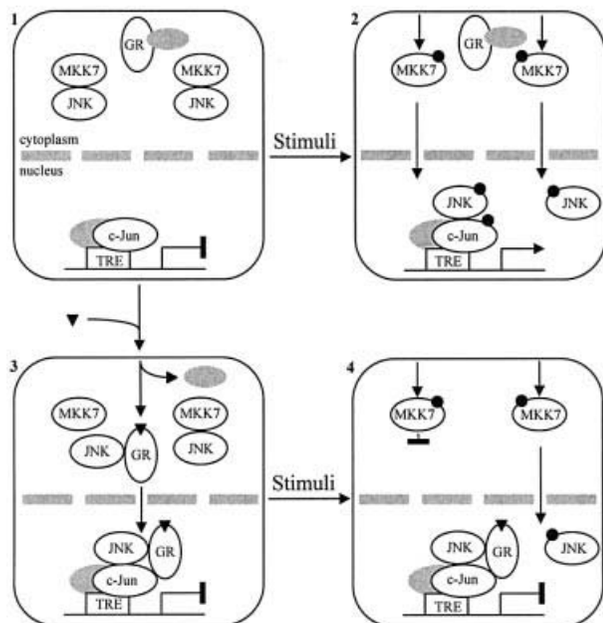


Fig. 7. GC action on the JNK pathway. (1) In the absence of GCs, GR is located in the cytoplasmic compartment and associated with multiprotein complexes. Likewise, in non-stimulated cells, JNK associated with MKK7 is tied by scaffold proteins into specific signaling modules. In the nuclear compartment, AP-1 complexes containing c-Jun are bound to the TPA response elements (TREs) found in the regulatory sequences of the AP-1 target genes, although the AP-1-dependent transcription is turned off. (2) Upon stimulation, the signal is transduced along the JNK pathway signaling modules by a cascade of phosphorylation events (filled circles) that ends in the activation of JNK. Active JNK dissociates from the signaling module and translocates into the nuclear compartment where it associates with and phosphorylates the c-Jun N-terminal domain. Thereafter, c-Jun phosphorylation induces the dissociation of a repressor complex resulting in AP-1 activation and, in consequence, triggers transcription of AP-1-responsive genes. (3) In the presence of GCs (inverted triangles), hormone-bound GR dissociates from the multiprotein complexes and exposes a JNK docking site. Some JNK molecules dissociate from the signaling modules, bind to GR and travel together with GR into the nuclear compartment where they associate with c-Jun. In these conditions, upon stimulation, JNK-deficient signaling modules fail to transduce the signal, resulting in the GC-induced inhibition of JNK pathway activation. In consequence, fewer molecules of active JNK are produced and, hence, enter into the nucleus. In addition, these active JNK molecules have to compete for the c-Jun docking sites with the already c-Jun-bound inactive JNK molecules.

GCs inhibit SEK1-dependent activation of JNK (Caelles *et al.*, 1997).

Here we demonstrate that GR binding to JNK is mediated by docking interactions analogous to those described to underlie other relevant traits of MAPK signaling pathways, such as sequential and specific activation and inactivation, substrate recognition and subcellular location of MAPKs (Tanoue and Nishida, 2002). In the LBD of GR, we have identified an amino acid motif that greatly resembles a D-box (Jacobs *et al.*, 1999), the type of MAPK docking site found in JNK-interacting proteins such as c-Jun or JIP-1. Within this amino acid sequence, GR, c-Jun and JIP-1 display identical or highly conserved residues in positions that are critical for binding to JNK (Kallunki *et al.*, 1996; Bonny *et al.*, 2001; Barr *et al.*, 2002; Sprowles and Wisdom, 2003). Consistently, the GR-LBD is sufficient to mediate binding to JNK and,

most significantly, this binding is competed specifically *in vivo* by a short peptide already validated as a JNK docking site, JNKI1 (Bonny *et al.*, 2001). Functionally, we show that this region of GR inhibits the JNK-induced transcription of an AP-1-dependent reporter in a similar way to the c-Jun N-terminal domain, thus further supporting its interaction with JNK *in vivo*. Remarkably, the amino acid sequence corresponding to this JNK docking site is highly conserved along GRs from a number of species, suggesting that the ability of this receptor to bind to JNK has been conserved during evolution. On the contrary, complete conservation of this docking site is not found in any other member of the NR superfamily, not even in its closest relative MR, suggesting that this may be a particular attribute of GR. In this regard, the inhibitory action on the JNK pathway of other NRs, such as the retinoic acid receptor, may be restricted to alternative mechanisms such as the induction of MKP-1 expression (Lee *et al.*, 1999; Xu *et al.*, 2002). It remains an open question as to whether the specific ability of GR to bind to JNK is related to its higher capacity, compared with other NRs, to inhibit the JNK pathway and transrepress AP-1. Taking advantage of the non-conservative amino acid exchanges, with respect to GR, in this region of the MR, we have disrupted the basic submotif of the JNK docking site of GR without significantly affecting hormone binding and shown that the single substitution of Arg576 by asparagine, the amino acid found in the same position in human MR, or by an acidic residue such as aspartate abolishes hormone-induced binding to and inhibition of JNK. Experiments are in progress to test if the poor AP-1 transrepression activity of MR (Pearce and Yamamoto, 1993; Heck *et al.*, 1994) might be improved by exchanging the amino acids of MR which are different from GR within this region, with the aim of providing MR with a JNK docking site and, eventually, with the ability to bind to and inhibit JNK.

Using several mutants of GR, such as GR-LBD, GR^{dim}, GR^{NLI}, GR^{R576N} and GR^{R576D}, as well as GC analogs such as RU486, we show that GR binding to JNK mediates two actions of GCs: inhibition of JNK pathway activation and induction of inactive JNK nuclear transfer. Dissociation of these two hormone actions demonstrated that nuclear accumulation of JNK is not required for downregulation of the JNK pathway. Therefore, we conclude that JNK activation is inhibited simply by its binding to the docking site offered by hormone-bound GR, in a similar manner to the JIP-1-based JNK inhibitor peptides (Bonny *et al.*, 2001; Barr *et al.*, 2002). Moreover, the results obtained with GR mutants with a constitutive cytoplasmic location indicate that binding, and hence inhibition, occurs in the cytoplasm. Additionally, by binding to GR, JNK may be provided with a nuclear localization signal and hence would be transferred to the nuclear compartment together with GR. Furthermore, we show that receptor dimerization is not required for binding to JNK, further confirming the association of JNK pathway inhibition with the transrepressive and DNA binding-independent function of GR.

We propose that once inside the nucleus via the action of GCs, inactive JNK is recruited to AP-1-bound response elements, such as those in the *c-jun* gene. Moreover, we show that the GR agonist Dex increases loading of both

GR and JNK onto these AP-1 response elements, while the GR antagonist RU486 is less effective in inducing GR and completely fails to induce JNK to associate with these AP-1 sites. Significantly, the failure of RU486 in binding to, inhibiting and inducing nuclear transfer of JNK is consistent with its weak AP-1 transrepressive activity (Vayssière *et al.*, 1997; Rogatsky *et al.*, 2001). Inactive JNK bound to the c-Jun N-terminal domain may block further interaction with and phosphorylation by activated JNK and thereby keep AP-1-dependent transcription repressed (Weiss *et al.*, 2003).

The negative regulation of MAPK pathways by GCs is achieved by alternative non-exclusive mechanisms and, most importantly, appears to mediate relevant physiological and pharmacological actions of GCs. Our data reveal a novel mechanism by which GCs target the JNK cascade, a major MAPK signal transduction pathway involved in AP-1 activation. Therefore, our results give further support to and rationale for pharmacological intervention in MAPK pathways.

Materials and methods

Plasmid constructs

To construct pSG424-Gal4-GR(540–738), the *Pst*I–*Hind*III DNA fragment from rat GR encoding amino acids 540–738 was cloned into pBS, excised with *Bam*HI–*Kpn*I and inserted into pSG424. To construct the plasmids pcDNA3-GR, pCEFL-KZ-HA-GR, pEBG-GR, pCEFL-KZ-HA-GR-LBD and pEBG-GR-LBD, the nucleotide sequence of rat GR encoding amino acids 1–795 or 525–795 was PCR-amplified using the following primer pairs: 5'-AGCTGGATCCACCATGGACTCCAAAG-AATCCTTA-3'/5'-AGCTACGCGCGCTCCATTTTGTATGAAACA-GAAGCTT-3' and 5'-AGCTGGATCCACCATGGAGTCTCAACAAG-CACT-3'/5'-AGTACGCGCGCTCATTTTTGTATGAAACAGA-AGCTT-3', respectively. Restricted PCR fragments were cloned into pCEFL-KZ-HA and pEBG between the *Bam*HI and *Nor*I restriction sites. pMTG-myc-GR_{R576N} and pMTG-myc-GR_{R576D} were obtained with the QuikChange Site-Directed Mutagenesis kit (Stratagene) using the following primer pairs: 5'-GTTCAGATTACGCATGGAATATTATG-ACCACACTC-3'/5'-GAGTGTGGTCATAATATCCATGCTGAATC-TGGAAC-3' and 5'-CAGCATGGGACATTATGACCACACTCAAC-3'/5'-GTCATAATGTCATGCTGAATCTGGAAC-3', respectively. All constructs were analyzed by DNA sequencing. Other plasmids have been described elsewhere (Heck *et al.*, 1994; Caelles *et al.*, 1997; Savory *et al.*, 1999; Chadee *et al.*, 2002).

Cell culture and transfection

Cos-7 and HeLa cells were grown in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal calf serum (FCS). Cells were serum starved in DMEM plus 0.5% FCS for 16 h. Unless indicated, Dex (10^{-6} M) or vehicle (ethanol) was added for 45 min before the cells were harvested, or stimulated with TNF- α (10 ng/ml) or UV (30 J/m²) and harvested 20 min later. Cells seeded in 100 mm tissue culture plates were transiently transfected by lipofection following the manufacturer's recommendations (Invitrogen) and using the following amounts of expression vectors: 0.5 μ g of pEBG, pEBG-JNK, pCEFL-KZ-HA-JNK, pEBG-GR or pEBG-GR-LBD and 1 μ g of pcDNA3-GR, pSB-GR, pSB-GR(A458T), pMTG-myc-GR, pMTG-myc-GR_{NL1}, pMTG-myc-GR_{R576N}, pMTG-myc-GR_{R576D}, pCEFL-KZ-HA-GR pCEFL-KZ-HA-GR-LBD or pCM15-Flag-MKK7, as indicated. Cells were harvested 24 h after transfection. Gene reporter assays were performed as described (Caelles *et al.*, 1997).

GST pull-down and co-immunoprecipitation assays

Cells were suspended in low-salt buffer [20 mM HEPES pH 7.4, 2 mM EGTA and 2 mM MgCl₂, 1 mM phenylmethylsulfonyl fluoride (PMSF) and 20 μ g/ml aprotinin] and lysed mechanically. Immunoprecipitation of HA-tagged proteins and endogenous JNK and MKK7 was performed using the antibodies 12CA5 (BabCO), sc-474 and sc-7104 (Santa Cruz Biotechnology Inc.), respectively. For the GST pull-down assays, cell lysates were incubated with 20 μ l of glutathione–Sepharose beads. After

three washes with low-salt buffer, immunocomplexes or glutathione–Sepharose precipitates were subjected to SDS–PAGE and analyzed by immunoblotting.

Immunoblotting

JNK, GR, MKK7, GST-, Flag-, HA- and myc-tagged proteins were detected by immunoblotting using the antibodies sc-474, sc-8992, sc-7104 and sc-138 from Santa Cruz Biotechnology, Inc., M2 antibody from Kodak, 12CA5 from BabCO, and 9E10 (a kind gift from Dr J.Ayté), respectively. Immunoblots were performed according to the enhanced chemiluminescence (ECL) detection system from Amersham Pharmacia.

Immunofluorescence analysis

Serum-starved HeLa or transiently transfected Cos-7 cells were treated with Dex, RU486 or vehicle. Thereafter, cells were processed as described (González *et al.*, 2000). Primary antibodies against GR (sc-8992, Santa Cruz Biotechnology, Inc.), JNK (15701A, Pharmingen) and HA tag (BabCO, 12CA5) were used. Immunofluorescence was analyzed by laser scanning confocal microscopy, and nuclear fluorescence quantified by the Metamorph Imaging System (Universal Imaging Corporation).

ChIP assay

ChIP assays were performed in serum-starved HeLa cells (2.5×10^7 cells per condition) treated with vehicle or hormone (Dex or RU486, 10^{-7} M) and following the method described by Rogatsky *et al.* (2001). Immunoprecipitation was performed using 2.5 μ g of anti-JNK, anti-GR or anti-GST antibody (sc-474, sc-8992 or sc-138, respectively, from Santa Cruz Biotechnology, Inc.) Immunoprecipitated DNA was suspended in 50 μ l of $0.1 \times$ TE. Aliquots of 5 μ l of each sample were PCR-amplified in the presence of 2.5 μ Ci of [α -³²P]dCTP with either the primer pair 5'-CAAGGACGTCAGCCCACAATG-3'/5'-ACACTCAGTGCAACTCT-GAG-3', which flanks the two AP-1 response elements found in the 5'-regulatory region of the *c-jun* gene and gives rise to a 347 bp fragment, or the primer pair 5'-CCAGCGTATCTATATGGAAATTG-3'/5'-AAAGA-TGGCCTTTGTCTTA-3', which amplifies a 287 bp fragment corresponding to the 3' end of *c-jun*. One-fifth of each PCR was electrophoresed onto a 6% polyacrylamide gel in $0.5 \times$ TBE and radioactive labeling was quantified using a PhosphorImager.

JNK immunocomplex assay

JNK and HA-JNK were immunoprecipitated from cell extracts using anti-JNK (sc-474 from Santa Cruz Biotechnology) and anti-HA antibodies (12CA5 from BabCO), respectively, and the JNK activity associated with immunoprecipitates was determined and quantified as described (Caelles *et al.*, 1997).

Acknowledgements

We thank Drs A.C.B.Cato, R.J.G.Haché, J.Ayté and G.Gil for providing the plasmids and reagents used in this study, C.Vila for her excellent technical assistance, the Confocal microscopy service of the Scientific and Technical Services of the University of Barcelona for their help with the confocal analysis, and Tanya Yates for style correction. This work was supported by grants from the Plan Nacional de I+D+I (SAF2001-3347) from the Ministerio de Ciencia y Tecnología and the Fundació 'La Caixa' (99/032-00). A.B. and M.N. were supported by a fellowship from Ministerio de Educación y Cultura and a grant from the Generalitat de Catalunya, Spain, respectively.

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Received March 27, 2003; revised September 16, 2003;
accepted October 2, 2003